Abstract

Advances in treating peripheral nerve lesions have resulted from research in nerve regeneration and the use biomaterials as well as synthetic materials. When direct tensionless repair of peripheral nerve lesions is not possible, nerve conduits may be used to bridge digital sensory nerve gaps of ≤3 cm. Nerve autograft is the benchmark for larger, longer, mixed, or motor nerve defects. Biologic, autogenous conduits—typically veins or, rarely, arteries—have demonstrated their utility in nerve gaps <3 cm in length. Three types of bioabsorbable conduit have been approved by the US Food and Drug Administration, constructed of collagen, polyglycolic acid, or caprolactone. Caprolactone conduits have been found to be equivalent in results to autograft. Collagen conduits are next best, and polyglycolic acid conduits are functionally inferior. Further research and prospective, multicenter, large-scale trials are needed to help establish the role of synthetic, bioabsorbable conduits in peripheral nerve reconstruction.

Complex and technically demanding to manage, segmental nerve defects pose a challenge for even the most skilled surgeon. Tension-free repair of nerve lacerations is the optimal surgical treatment. When tensionless direct repair cannot be achieved, interposed nerve autograft is the benchmark. However, nerve autograft results in increased surgical time and donor site morbidity, thereby justifying the search for better options. Table 1 lists the current options for bridging nerve gaps. Nonneural, hollow, tubular interposition substitutes known as nerve conduits include autogenous vein or artery grafts and synthetic tubes. Sometimes these conduits are referred to as nerve guides. Although the use of acellular cadaver nerve allografts is also increasing, clinical studies are limited. Here, we review the use of hollow, tubular nerve conduits as a method for reconstructing nerve gaps.

History

The idea of repairing nerve gaps with hollow conduits, also known as tubulation, dates back to the late 1800s, when Gluck proposed using decalcified bone tubes for this purpose. In 1891, Bungner bridged a canine sciatic nerve gap with a segment of human brachial artery. Platt, in 1919, reported clinical application of 6-inch vein graft for radial (musculospiral) nerve reconstruction, with no functional return. Lundborg et al, in 1982, bridged rat sciatic nerves with silicone tubes, although silicone never became popular because of concerns of nerve constriction. Walton et al, in 1989, reported encouraging results in a retrospective...
A series of digital nerve injuries that had been reconstructed by vein conduits. In 1990, Chiu and Strauch reported a successful prospective series of autogenous vein nerve conduits (AVNCs) compared with nerve autografts for digital nerve gaps ≤3 cm in length. Mackinnon and Dellon bridged clinical nerve gaps of ≤3 cm with biodegradable polyglycolic acid (PGA) in 1990. Since the early 1990s, the number of nerve conduit studies has been steadily increasing. Table 2 summarizes the data from a number of these studies.

### The 3-cm Limit on Nerve Conduit Length

There is a generally accepted upper limit of 3 cm on nerve conduit length. Most reported series of nerve conduits for reconstruction of digital

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**Table 1**

<table>
<thead>
<tr>
<th>Conduit Type</th>
<th>Bridge</th>
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<tr>
<td>Nerve</td>
<td>Autograft, allograft</td>
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<tr>
<td>Biologic</td>
<td>Vein, artery</td>
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<tr>
<td>Synthetic</td>
<td>Collagen (NeuraGen, Integra LifeSciences, Plainsboro, NJ; Neuroflex and NeuroMatrix, Stryker Orthopaedics, Mahwah, NJ)</td>
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<td></td>
<td>Polyglycolic acid (NeuroTube, Synovis Micro Companies Alliance, Birmingham, AL)</td>
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<td></td>
<td>Caprolactone (Neurolac Nerve Guide, Polyganics BV, Groningen, The Netherlands)</td>
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**Table 2**

<table>
<thead>
<tr>
<th>Study</th>
<th>Conduit Type</th>
<th>Study Type (level of evidence)</th>
<th>Outcomes Measured</th>
<th>Conclusions</th>
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<tr>
<td>Chiu and Strauch²</td>
<td>AVNC versus nerve autograft</td>
<td>Prospective cohort (II)</td>
<td>Static, moving 2PD; patient satisfaction questionnaire</td>
<td>AVNC comparable to nerve autografts for gaps ≤3 cm</td>
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<tr>
<td>Flores⁶</td>
<td>AVNC</td>
<td>Case-control (III)</td>
<td>Static 2PD; SWM</td>
<td>AVNC of sural nerve biopsy defects did not shorten time to sensory recovery. Quality of reinnervation better than control subjects.</td>
</tr>
<tr>
<td>Rinker and Liau⁷</td>
<td>AVNC versus PGA (NeuroTube⁸)</td>
<td>Randomized controlled trial (II)</td>
<td>Static, moving 2PD</td>
<td>No difference in sensory results between groups for repair of nerve gaps of 4–25 mm. Similar cost profile for both groups. More complications in PGA group with two extrusions requiring reoperation, although not statistically significant.</td>
</tr>
<tr>
<td>Lohmeyer et al⁸</td>
<td>Collagen (NeuraGen⁹)</td>
<td>Prospective cohort (II)</td>
<td>Static 2PD</td>
<td>75% good to excellent results</td>
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<tr>
<td>Bertleff et al⁹</td>
<td>Caprolactone (Neurolac Nerve Guide²) versus primary repair</td>
<td>Multicenter, blinded, randomized controlled trial (II)</td>
<td>Static, moving 1PD and 2PD</td>
<td>Recovery of sensation as good as that of control subjects. Time for repair greater by 14 min, and more complications reported in experimental conduit group.</td>
</tr>
<tr>
<td>Weber et al¹⁰</td>
<td>PGA (NeuroTube⁹) versus primary repair (end-to-end or with nerve graft)</td>
<td>Multicenter, randomized, prospective controlled trial (II)</td>
<td>Moving 2PD</td>
<td>No statistically significant difference in groups in terms of overall results. Conduits superior to primary repair for gaps ≤4 mm, superior to nerve autograft for gaps &gt;8 mm.</td>
</tr>
</tbody>
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¹PD = 1-point discrimination, 2PD = 2-point discrimination, AVNC = autologous vein nerve conduit, PGA = polyglycolic acid, SWM = Semmes-Weinstein monofilament
⁸ Synovis Micro Companies Alliance, Birmingham, AL
⁹ Integra LifeSciences, Plainsboro, NJ
ⁱ⁰ Polyganics BV, Groningen, The Netherlands
nerve defects adhere to the 3-cm limit. Mackinnon11 recently reported that the indications for nerve conduits are limited to small-diameter, noncritical sensory nerves with a gap of <3 cm. Strauch et al.,12 in a rabbit peroneal nerve study that compared the results of axonal regeneration using vein conduits from 1 to 6 cm in length, found excellent growth and function ≤3 cm, with deteriorating results at lengths >3 cm.

Investigators have attempted to overcome the 3-cm limit by experimentally inserting Schwann cells13 or portions of nerve or muscle within the conduit vein or tube; however, this practice has not found widespread clinical use or acceptance. The author of one study used 5-cm vein conduits for sural nerve defects following nerve biopsy and reported successful results.6 However, the sensory recovery of the conduits in this study was compared with that of control subjects in which the entire sural nerve had been harvested. The control subjects also obtained satisfactory recovery of protective sensation within 10 months, thus questioning the validity of the model.

**Use of Conduits for Larger Diameter Nerves With Mixed Motor and Sensory Fibers**

Most clinical studies of nerve conduits have targeted digital sensory nerve defects. Conduit reconstruction of larger nerves, such as the median, ulnar, or radial, has not been as well studied. Moore et al14 recently reported on four patients with unsuccessful conduit repair of larger nerves, including median, ulnar, and brachial plexus nerves. Conversely, Donoghoe et al15 reported successful repair of 3-cm median nerve gaps using PGA conduits in cable formation. Stanec and Stanec16 bridged a 2.9-cm ulnar nerve gap with an expanded polytetrafluoroethylene tube in 1998. While anecdotal reports of larger or mixed nerve gap reconstruction with conduits have appeared, there is insufficient clinical support for the routine use of nerve conduits over nerve autografts for this indication. Although synthetic conduits are fabricated in wider diameters, this does not imply successful outcomes when they are used for larger nerves. Prospective randomized clinical trials are needed to evaluate the role of nerve conduits for mixed or purely motor nerve defects, and/or for defects ≥3 cm in length.

**Autogenous Conduits**

Autogenous conduits are usually veins (AVNCs) or, rarely, arteries. Kosutic et al17 published a case series of two homolateral digital arteries used to bridge 2- and 3-cm digital nerve defects that, at 2-year follow-up, demonstrated improved static two-point discrimination to protective levels in both patients (≤7 mm). The technique for AVNC involves resection back to healthy nerve, harvesting a vein twice the diameter of the nerve and 50% longer than the gap, reversing the polarity of the vein, and intussuscepting the nerve ends into the vein lumen with microsutures18 (Figure 1). Numerous clinical reports of AVNC have borne out its utility in nerve gaps <3 cm in length. Chiu and Strauch,2 in a prospective study of 22 patients with painful neuromas or segmental nerve injuries of <3 cm, found that AVNCs produced clinical results similar to those of sural digital nerve grafts but inferior to those of primary end-to-end repair. The authors did not inject saline solution or heparin into the vein graft.

A recent prospective randomized clinical trial that compared AVNCs to PGA conduits for digital nerve gaps from 4 to 25 mm found equivalent sensory results between the groups; however, there were more complications in the PGA group, including two extrusions requiring re-operation.7 The theoretic concern of vein graft collapse, that the vein tube will flatten and block nerve regeneration, has not been borne out clinically, and there is no clear evidence that inserting muscle or other material into the vein is superior to no interposition.

**Synthetic Conduits**

Three types of bioabsorbable conduits are currently approved by the US Food and Drug Administration (FDA) for use, constructed of collagen, PGA, or caprolactone. Insertion of nerve conduits requires isolation of the defect, followed by selection of the diameter and length of the tube. The technique for inserting the nerves ends into the tubes is generally as follows: The suture needle is

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*Figure 1*

placed from outside the tube into the lumen. An epineural suture is placed in the nerve, and the suture is passed from inside the tube to outside the tube, thereby pulling the nerve end into the tube. Finally, the suture is tied over the conduit (Figure 2).

Shin et al. compared the performance of nerve autograft to that of caprolactone, collagen, and PGA conduits in a rat sciatic nerve model with a 10-mm defect. Caprolactone conduits were found to be equivalent to autograft; collagen conduits performed next best; and PGA conduits produced greatly inferior functional results and had structurally completely collapsed by 12 weeks.

**Collagen**

Types I and III collagen make up 49% of peripheral nerve proteins, with type I most predominant. Type I collagen is biocompatible and constitutes most conduits. The semi-permeable nature of collagen conduits promotes diffusion and resorption by 9 months. NeuraGen (Integra LifeSciences, Plainsboro, NJ), NeuroMatrix collagen matrix (Stryker Orthopaedics, Mahwah, NJ), and Neuroflex collagen matrix (Stryker) nerve cuffs are examples of commercially available collagen conduits. Figure 3 shows intraoperative photographs demonstrating the nerve conduit in situ used for repair of a common digital nerve.

Bushnell et al. reported a 2-year follow-up of a level IV case series of 12 collagen conduit repairs of digital nerve gaps ranging in length from 10 to 12 mm. American Society for Surgery of the Hand guidelines with static two-point discrimination, Disabilities of the Arm, Shoulder, and Hand (DASH) scores, and Semmes-Weinstein testing were used to measure outcome. Of the nine patients available for follow-up, four (44%) had excellent results, four had good results (44%), and one had a fair result (11%), with average DASH score of 10. Lohmeyer et al. performed a prospective cohort study involving collagen conduits to repair 12 digital nerves with an average 12.7-mm gap. One-year follow-up demonstrated 33% excellent sensory recovery and 42% good sensation, with 8% poor sensation and 8% no sensory recovery. Currently there are no randomized controlled trials examining collagen tubes. Additionally, grading of outcomes by using two-
point discrimination is not standardized. No studies have examined motor recovery with collagen tubes.20

**Polyglycolic Acid**

Early synthetic conduit research was performed using PGA. This tube is regarded as more flexible and porous than others, thereby allowing diffusion to aid in regeneration, with resorption occurring in 6 months.20 Mackinnon and Dellon,4 in a prospective level IV case series, examined 15 patients undergoing secondary nerve reconstructions with PGA tubes of digital nerve gaps measuring approximately 17 mm. These authors found that 33% of patients had excellent sensory recovery, 53% good recovery, and 14% poor recovery. Sensory data was gathered using the British Medical Research Council sensory grading scale with moving and static two-point discrimination. Excellent recovery was defined as static two-point discrimination. Experiments were superior to primary repair for nerve gaps of ≤4 mm (a gap length usually amenable to primary repair) and superior to nerve autograft for gaps ≥8 mm. The study design, however, included a wide range of variables that likely prohibit drawing the conclusion that conduit repair is superior to primary repair or autograft; primary repair or autograft is still considered by most to be superior to conduit repair.

**Caprolactone**

An aliphatic polyester, poly(DL-lactide-caprolactone), was first demonstrated in rat models to bridge 10-mm sciatic nerve gaps, with complete degradation in 1 year.20 Further research has raised the issues of conduit inflexibility and unabsorbed fragments. Bertleff et al9 performed a multicenter blinded randomized controlled trial of 30 patients with 34 nerve injuries using Neurolac nerve tubes (Polyganics BV, Groningen, The Netherlands), which are made of caprolactone, compared with primary repair for digital nerve lacerations. Gaps of 6 to 8 mm were repaired with Neurolac tubes. Digital nerves without gaps were repaired primarily with 8-0 or 9-0 nonabsorbable suture. Pressure sensation and two-point discrimination were evaluated using a noninvasive, computer-assisted force transducer. Moving and static two-point discrimination was 7 to 10 mm for both the experimental and control groups. Time for repair was greater in the conduit group by 14 minutes; complications were greater in the Neurolac group.

This study represents preliminary evidence that caprolactone nerve tubes produce results comparable to those of primary digital nerve repair, although additional studies are needed.

**Summary**

Research in nerve regeneration and biomaterials has led to advancements in managing peripheral nerve lesions. When a direct tensionless repair is not possible, conduits may be used to bridge digital sensory nerve gaps of ≤3 cm, with nerve autograft remaining the benchmark for larger, longer, mixed, or motor nerve defects. Biologic, autogenous conduits have demonstrated their utility in nerve gaps of ≤3 cm. Bioabsorbable conduits of collagen, PGA, and caprolactone have been approved by the US FDA; caprolactone conduits have been found to be equivalent in results to autograft. As tissue bioengineering advances provide ways to enhance growth and increase neurotropism, further research may expand the indications for use of nerve autografts, autogenous conduits, and synthetic conduits. There is clearly a need for prospective, multicenter, large-scale trials to aid in surgical decision making in the future of peripheral nerve reconstruction.

**References**

References printed in bold type are those published within the past 5 years.


3. Lundborg G, Gelberman RH, Longo FM, Powell HC, Varon S: In vivo regeneration of cut nerves encased in


